

COMMUNICATION

[View Article Online](#)
[View Journal](#) | [View Issue](#)



Cite this: *Nanoscale Adv.*, 2019, **1**, 4669

Received 20th August 2019
Accepted 12th October 2019

DOI: 10.1039/c9na00521h
rsc.li/nanoscale-advances

The behavior of Li/Na-ion capacitors (LICs/NICs) is largely limited by the low number of electroactive sites in conventional insertion-type anodes. In this work, we demonstrated a novel doping-vacancy double-defective and conversion-type Mn–O–F ultrafine nanowire (denoted as $\text{MnF}_2\text{-E}$) anode to boost the number of electroactive sites for enhanced LICs/NICs. Owing to the unique hetero oxygen-doping and intrinsic fluorine-vacancy double defects, the Mn–O–F nanowires exhibited superior electroactive sites and thus dramatically enhanced Li/Na-ion storage capability than pristine MnF_2 micro/nano-crystals. Both the optimal MnF_2 screened by orthogonal experiments and derived Mn–O–F anodes and commercial activated carbon (AC) cathode were used to construct $\text{MnF}_2//\text{AC}$ and $\text{MnF}_2\text{-E}/\text{AC}$ LICs/NICs, which were optimized by tuning the active mass ratios of the cathode/anode and the working voltage windows of the hybrid capacitors. The LICs/NICs based on the Mn–O–F anode demonstrated a considerably superior performance than the devices based on the MnF_2 anode under the optimal voltages of 0–4 V and 0–4.3 V. The Mn–O–F anode exhibited dominant diffusion/surface-controlled kinetics for Li/Na-ion storage, respectively, showing a major conversion mechanism for the charge storage processes. This work provides a new concept of double-defective and conversion-type electrode materials to improve the Li/Na-ion storage capability and will have a significant impact on the relevant fields.

Introduction

Energy is increasingly inseparable from human life with the development of society; however, the disadvantages of the depletion of fossil fuel resources and environmental pollution limit their applications. Thus, electrochemical energy storage has become one of the most important strategies to solve these

Key Laboratory of Environmentally Friendly Chemistry and Applications of Ministry of Education, College of Chemistry, Xiangtan University, Xiangtan, Hunan 411105, P. R. China. E-mail: drm8122@163.com; drm8122@xtu.edu.cn

† Electronic supplementary information (ESI) available. See DOI: [10.1039/c9na00521h](https://doi.org/10.1039/c9na00521h)

Engineering doping-vacancy double defects and insights into the conversion mechanisms of an Mn–O–F ultrafine nanowire anode for enhanced Li/Na-ion storage and hybrid capacitors†

Yongfa Huang, Rui Ding,  * Danfeng Ying, Wei Shi, Yuxi Huang, Caini Tan, Xiujuan Sun, Ping Gao and Enhui Liu

issues.¹ Electrochemical energy storage, which is necessary for future large-scale power grids, is sustainable, green and responsive and can be used in electrochemical engines and smart devices.² Electrochemical capacitors (ECs) or supercapacitors (SCs) as an important part of electrochemical energy storage have attracted wide interest from researchers,³ especially lithium-ion capacitors (LICs)^{4,5} and sodium-ion capacitors (NICs).^{6,7} Furthermore, non-aqueous hybrid ion capacitors^{8,9} usually show a higher working voltage, larger energy density and wider temperature scope than aqueous hybrid ion capacitors.¹⁰

LIC was proposed by Amatucci's team in 2001.^{11,12} It uses a battery-type material as the negative electrode with high energy density and an electric double layer (EDL) carbon material as the positive electrode with high power density; thus, LICs can export high energy and power density simultaneously. However, the performance of LICs is largely limited by anode materials, which have low electrochemical activity. Thus, finding new anodes with high electroactive sites has become an important research direction. Various advanced materials such as Nb_2O_5 ,¹³ MnO ,¹⁴ and BiVO_4 ¹⁵ have been reported as intriguing anodes for LICs.

Nowadays, the depletion of lithium resources has caused widespread concern and more seriously, the development of lithium resources cannot keep up with the development of lithium-ion energy storage devices.¹⁶ Therefore, it is very important to find electrochemical energy storage devices with low cost, high safety, high energy/power densities and long cycle stability. It is particularly attractive to develop other ion energy storage devices for this purpose. Sodium (Na) ions are highly abundant and low cost in nature and have received great attention from researchers as alternatives to lithium ions.¹⁷ Sodium ion capacitors (NICs) also have the advantages of sodium ion batteries of high energy density and EDL capacitors of high power density. Some NICs even exhibit the same performance as LICs, such as $\text{Nb}_2\text{O}_5@\text{C}$ /reduced graphene oxide (rGO),¹⁸ TiO_2/C ,¹⁹ N-TiO_2 ,²⁰ 3D framework carbon (3DFC),²¹ peanut skin-derived carbon nanosheets-A (SCN-A),²²

Electrode materials play a key role in electrochemical energy storage. The choice of materials should be based specifically on the following reasons: firstly, high specific surface area can afford rich electroactive sites, which can ensure high specific capacity. Secondly, high porosity can provide facile pathways for ion transportation and superior endurance for volume change, which can ensure high rate capability and stability. Thirdly, special morphologies such as nanowires and nanotubes, can facilitate the electron/ion transfer rates, which can further enhance the electrochemical performance.

In recent years, heteroatom doping has been widely used in electrochemical energy storage devices, which can greatly improve the surface properties of electrode materials, shorten the ion transport channels, and thus significantly improve the charge storage capability,^{23,24} which include (N or S)-doped graphene,²⁵ (B, N or S)-doped hard carbon,²⁶ and S-doped V_6O_{13-x} (VOS).²⁷ Furthermore, vacancy defects can increase the number of electroactive sites and enhance the electron/ion transfer kinetics of electrode materials,^{23,24} such as O-vacancy- Co_3O_4 ,²⁸ and Co-vacancy-CoSe₂.²⁹ Considering the above advantages, the Li/Na-ion storage capability of the pristine MnF₂ anode can be significantly increased by simultaneously introducing hetero oxygen-doping and intrinsic fluorine anion-vacancies.

In this work, we report hetero oxygen-doping and fluorine anion-vacancy double-defective Mn–O–F ultrafine nanowires as novel promising anodes for LICs and NICs. Due to the disadvantages of large particle size and low specific surface area of pristine MnF₂, we propose a simple method to solve these problems *via* the etching treatment of pristine MnF₂ micro/nanocrystals with NaBH₄ agents, which generates double-defective Mn–O–F ultrafine nanowires with high porosity and large specific surface area (the materials are denoted as MnF₂-E). The performance of MnF₂ candidates was firstly optimized *via* an orthogonal experiment (L_9).³⁴ The optimal MnF₂ (8#) and etched MnF₂ (8#)-E (Mn–O–F) anodes and commercial activated carbon (AC) cathode were used to construct the MnF₂/AC and MnF₂-E//AC LICs/NICs, which were optimized by tuning the active mass ratios of positive and negative electrodes and working voltage windows of the capacitors. Both the LICs and NICs based on the double-defective Mn–O–F anode showed much better electrochemical performance than the devices based on the MnF₂ anode. The kinetics and mechanisms of the MnF₂ and Mn–O–F anodes for both Li-ion and Na-ion storage were also investigated. Overall, this work addresses new insight into heteroatom doping and anion vacancy double-defective Mn–O–F anodes for advanced LICs and NICs and will have a significant impact on the development of advanced electrode materials for high-performance electrochemical energy storage devices.

Results and discussion

Physicochemical property

The optimal MnF₂ 8# (herein, the choice of MnF₂ 8# is owing to its overall superior specific capacity, rate capability and cycling behavior, Fig. S6 and Table S4, ESI[†]) and MnF₂ 8#-E (Mn–O–F)

materials were characterized by X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR), X-ray photoelectron spectroscopy (XPS), scanning electron microscopy (SEM), transmission electron microscopy (TEM), high-resolution TEM (HRTEM), selected area electron diffraction (SAED), X-ray energy dispersive spectroscopy (EDS), mapping, and nitrogen isothermal sorption with Brunauer–Emmett–Teller (BET) and Barrett–Joyner–Halenda (BJH) methods. Fig. 1A shows the XRD patterns of the MnF₂ 8# and MnF₂ 8#-E samples, which correspond to tetragonal MnF₂ with the space group of $P4_2/mnm$ (the crystal structures of MnF₂ 8# and MnF₂ 8#-E are shown in Fig. S1[†]). The XRD patterns of the other eight samples from the orthogonal experiment (Fig. S2 and Tables S1, S2[†]) also correspond to the tetragonal MnF₂ standard card, indicating the successful synthesis of the materials. It is apparent that the peak intensity of MnF₂ 8#-E is significantly weaker than that of MnF₂ 8#, which indicates that the particle size or crystallinity of the MnF₂ 8#-E sample decreased significantly. More interestingly, in the enlarged view of the crystal plane of (110), the peak position of MnF₂ 8#-E shows a slight negative shift by 0.03°, which may be due to oxygen heteroatom doping. The IR spectra of MnF₂ 8#-E and MnF₂ 8# in Fig. 1B demonstrate the presence of the Mn–O bond in MnF₂ 8#-E, which is direct evidence of the oxygen heteroatom doping. Fig. 1C shows the XPS survey scan spectra of MnF₂ 8# and MnF₂ 8#-E. The typical Mn 2p and F 1s can be clearly seen for MnF₂. Notably, the O 1s peak of MnF₂ 8#-E is significantly stronger than that of MnF₂ 8#, whereas the F 1s peak of MnF₂ 8#-E is obviously weaker than that of MnF₂ 8#, which is attributed to the fact that the oxygen heteroatoms largely occupy the positions of the fluorine atoms, resulting in the presence of rich fluorine vacancies owing to the charge

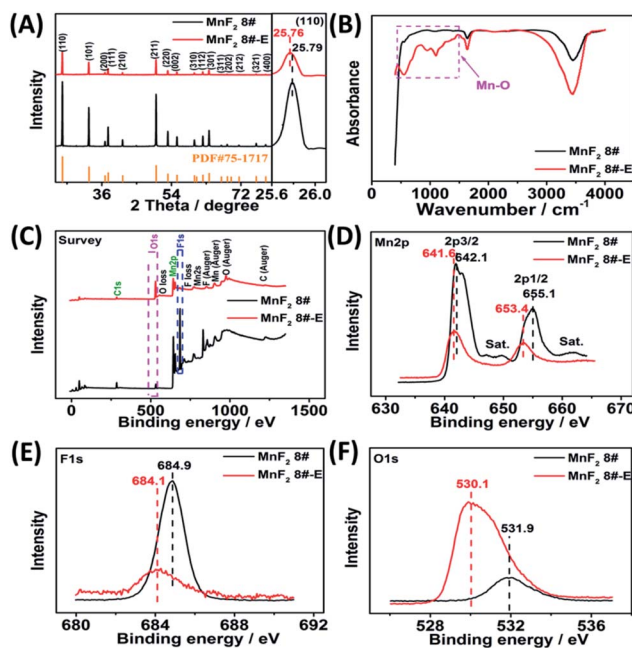


Fig. 1 XRD patterns (A), FTIR spectra (B), XPS survey scan (C), and XPS Mn 2p (D), F 1s (E) and O 1s (F) spectra of the MnF₂ 8# and MnF₂ 8#-E samples.



balance. The Mn 2p spectra of MnF_2 8# and MnF_2 8#-E are shown in Fig. 1D, where Mn 2p_{3/2} and Mn 2p_{1/2} are negatively shifted by 0.6 eV and 1.7 eV compared with the Mn 2p spectra of MnF_2 8#, respectively, which is due to the formation of the Mn–O bond by oxygen doping. The F 1s spectrum of MnF_2 8#-E in Fig. 1E is significantly lower than that of MnF_2 8#, accompanied by a negative shift of 0.8 eV for MnF_2 8#-E, which is due to the hetero oxygen doping and fluorine ion vacancy. Fig. 1F shows the O 1s spectra, where the O 1s peak of MnF_2 8#-E exhibits a negative shift of 1.8 eV compared to that of MnF_2 8#, which is due to the formation of a strong Mn–O bond by the hetero oxygen doping. Moreover, the ratio of Mn/O/F was estimated to be 1 : 0.934/0.132 based on the XPS results, and thus the formula of Mn–O–F can be expressed as $\text{MnO}_{0.934}\text{F}_{0.132}$. Herein, the doping of oxygen heteroatoms and fluorine vacancy double defects gave Mn–O–F a distinct advantage in electrochemical performance.

The SEM and TEM images of MnF_2 8# are shown in Fig. 2A and B, respectively, showing a size range of around 0.5–2 μm . The HRTEM image of MnF_2 8# in Fig. 2C shows the lattice fringes of 0.345 nm, which is matched with the (110) crystal plane of tetragonal MnF_2 . The SAED patterns in Fig. 2D exhibit the single-crystalline diffraction characteristics of MnF_2 8# with the marked standard crystalline planes of (110), which are in

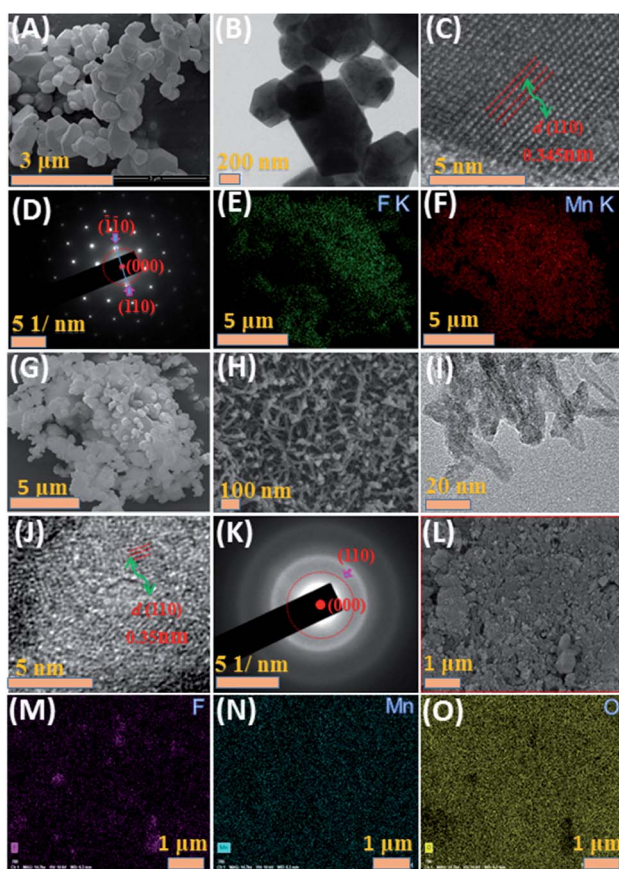
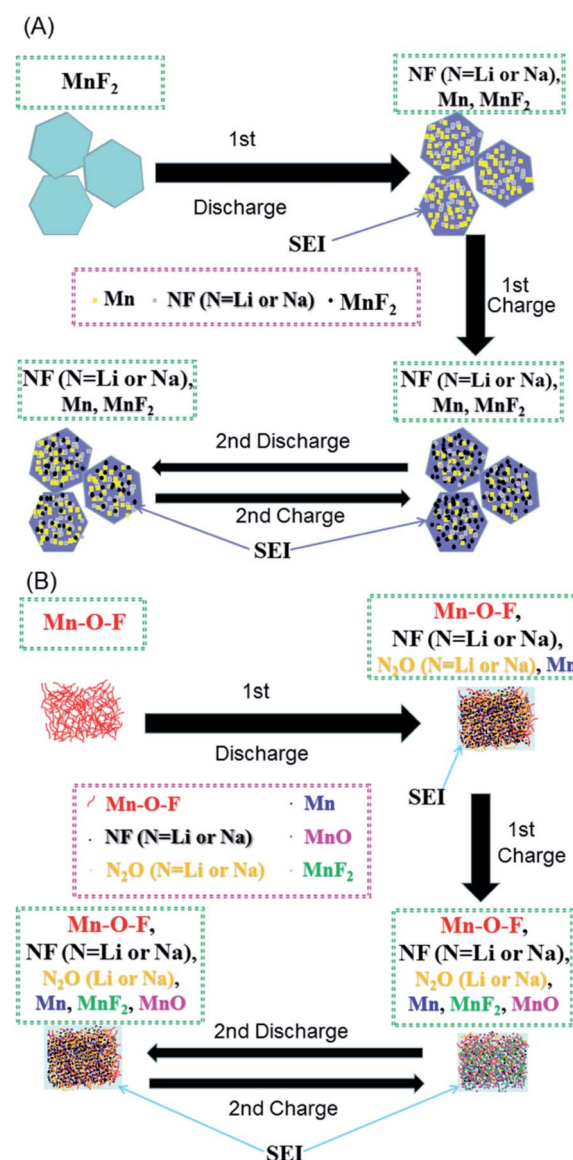


Fig. 2 SEM (A), TEM (B), HRTEM (C), SAED (D) and mapping (E–G) images of MnF_2 8# sample. SEM (H), TEM (I), HRTEM (J), SAED (K) and mapping (L–O) images of MnF_2 8#-E sample.

good agreement with the HRTEM and XRD results. Furthermore, the mapping images of MnF_2 8# in Fig. 2E–G demonstrate the uniform distributions of F/Mn species. Fig. 2H, I and S3[†] show the SEM/TEM images of MnF_2 8#-E, which exhibit an ultrafine nanowire morphology with a diameter of about 10 nm, and a much more porous structure can be clearly seen for MnF_2 8#-E (the possible formation process can be seen in Scheme 1, ESI[†]), which endows the MnF_2 8#-E sample with more electroactive sites than the pristine MnF_2 8# sample. The ultrafine nanowire morphology and porous structure also ensure a shorter ion transport length and provide a larger electrode/electrolyte interface for charge transport reactions, which is also reflected in its superior electrochemical performance.³⁰ The HRTEM image and SAED pattern of MnF_2 8#-E are shown in Fig. 2J and K, respectively, in which a typical amorphous



Scheme 1 Schematics of the possible reaction mechanisms for the MnF_2 (A) and MnF_2 -E (B) electrodes during the first two cycles of discharging/charging processes.

structure with a lightly larger lattice fringe of 0.35 nm corresponding to the (110) crystal plane can be detected. The mapping diagrams of MnF_2 8#-E in Fig. 2L-O demonstrate the even distribution of Mn, O, and F elements in the MnF_2 8#-E sample, which again prove hetero oxygen doping in the MnF_2 structure. The different colors of MnF_2 8# and MnF_2 8#-E in Fig. S4[†] and the element results from the EDS data in Fig. S5[†] also reflect the doping of oxygen heteroatoms and the change in the intrinsic structure of the materials. Further changes can be also seen from the nitrogen isothermal sorption data, pore volume and pore size distribution of MnF_2 8# and MnF_2 8#-E (Fig. 3A-F). Specifically, MnF_2 8# has a very small specific surface area of only $2.75 \text{ m}^2 \text{ g}^{-1}$, while MnF_2 8#-E exhibits a staggering value of $198.96 \text{ m}^2 \text{ g}^{-1}$, and the pore volume of MnF_2 8# is also very small ($0.008 \text{ cm}^3 \text{ g}^{-1}$) in comparison with the very large pore volume of MnF_2 8#-E ($0.95 \text{ cm}^3 \text{ g}^{-1}$), and the pore size of MnF_2 8# is mainly distributed at 2.23 nm in comparison with the bigger pore size distribution of MnF_2 8#-E (11.26 nm), which indicate an enormous increase in specific surface area, pore volume and pore size by the introduction of hetero oxygen doping and fluorine vacancy double defects, ensuring the exceptionally superior electrochemical performance of the double-defective Mn-O-F ultrafine nanowires for Li/Na-ion storage.

Performance for Li-ion storage

To evaluate the electrochemical performance of the MnF_2 (1#-9#) samples and select the optimal candidate from the orthogonal experiments, Li// MnF_2 half cells were assembled and

tested *via* cyclic voltammetry (CV) and galvanostatic charge/discharge (GCD) between 0.01 and 3 V (vs. Li/Li⁺). Based on the results of the orthogonal analysis and performance comparison in Tables S3, S4 and Fig. S6,[†] MnF_2 8# exhibited an overall superior performance. The cyclic voltammogram of the first three revolutions at 0.1 mV s^{-1} of MnF_2 8# is shown in Fig. 4A, with one minor peak located at about 0.96 V and a sharp peak at around 0.29 V for the first cathode scan, which coincide with the platforms and the sloped regions at the low voltage appearing in the original discharge branch of the GCD curves at 0.1 A g^{-1} (Fig. 4B), respectively. These peaks are largely owing to the conversion of MnF_2 into Mn/LiF and the formation of solid electrolyte interphase (SEI) films accompanied with a highly reversible interfacial Li-ion intercalation reaction within the Mn/LiF matrix in the low potential region, respectively.³¹⁻³⁵ Also, the minor anodic peak at 1.01 V can be ascribed to the partial conversion of Mn/LiF into MnF_2 .^{31,32} However, the redox peaks

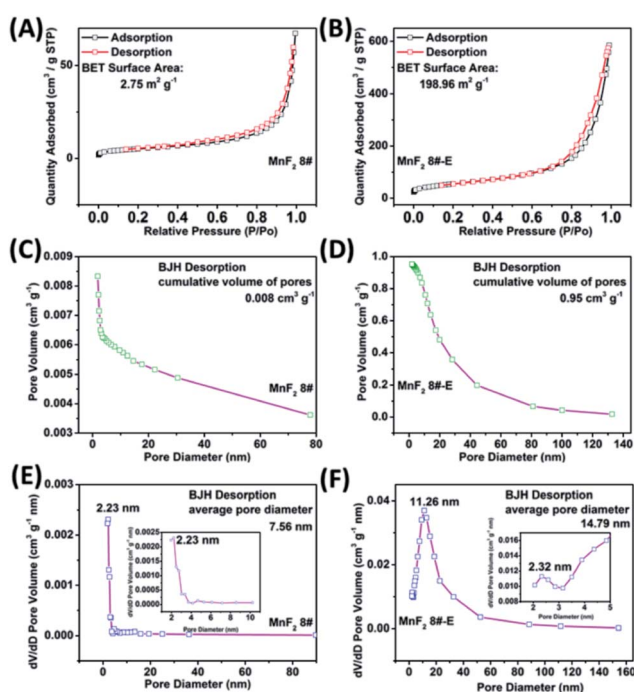


Fig. 3 N_2 isothermal sorption (A and B), pore volume (C and D) and pore size distribution (E and F) of the MnF_2 8# and MnF_2 8#-E samples, respectively.

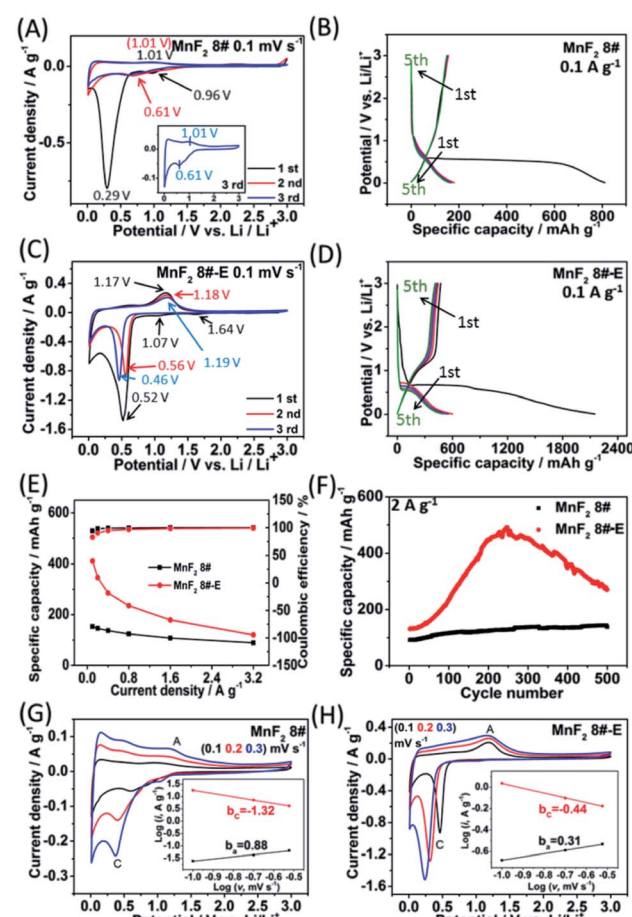


Fig. 4 CV plots for the first three cycles at 0.1 mV s^{-1} (A) and GCD curves for the 5th cycle at 0.1 A g^{-1} (B) of MnF_2 8# electrode, CV plots for the first three cycles at 0.1 mV s^{-1} (C) and GCD curves for the 5th cycle at 0.1 A g^{-1} (D) of MnF_2 8#-E electrode, specific capacity and coulombic efficiency derived from the respective 5th cycle at $0.1-3.2 \text{ A g}^{-1}$ of MnF_2 8# and MnF_2 8#-E electrodes (E), cycling stability for 500 cycles at 2 A g^{-1} of MnF_2 8# and MnF_2 8#-E electrodes (F), and CV plots for the 3rd cycle at $0.1, 0.2$, and 0.3 mV s^{-1} of MnF_2 8# and MnF_2 8#-E (insets show the plots of $\lg i$ vs. $\lg v$) (G and H).



were maintained at about 1.01/0.61 V in the subsequent cycles, and it can be seen from the charge–discharge curves that the capacity is basically unchanged in the subsequent cycles except for the much larger specific capacity of the original discharge, indicating that the internal structure is protected after the formation of the SEI films, and shows good electrochemical stability.^{31,32} Similar changing trends in the redox peaks and charge–discharge platforms can be also found from the CV curves and GCD plots of the other MnF_2 samples, as shown in Fig. S7 and S8.[†]

Fig. 4C shows the CV plots of MnF_2 8#-E at a sweep speed of 0.1 mV s⁻¹, with two weak cathode peaks at 1.64 V and 1.07 V and a sharp cathode peak at 0.52 V in the first cathode sweep, which may be due to the conversion of $\text{Mn}-\text{O}-\text{F}$ into $\text{Mn}/\text{LiF}/\text{Li}_2\text{O}$ and the formation of the SEI films, and the minor anodic peak at 1.17 V can be ascribed to the partial conversion of $\text{Mn}/\text{LiF}/\text{Li}_2\text{O}$ into MnF_2/MnO .^{14,31,32} In the next two cycles, the cathode peaks shifted slightly from 0.56 V to 0.46 V, and the anode peaks shifted slightly from 1.18 V to 1.19 V, indicating good reversibility. Fig. 4E and F show the specific capacity and cycle performance of the MnF_2 8# and MnF_2 8#-E electrodes, respectively, where MnF_2 8#-E provided the much bigger specific capacity of 411–120 mA h g⁻¹ at 0.1–3.2 A g⁻¹ in comparison with that of 153–89 mA h g⁻¹ at 0.1–3.2 A g⁻¹ for the MnF_2 8# electrode. The more obvious gap is also reflected in the cycle performance after 500 cycles at 2 A g⁻¹, where MnF_2 8#-E exhibited 271 mA h g⁻¹ (207% retention), whereas MnF_2 8# exhibited only 138 mA h g⁻¹ (152% retention). Moreover, the performance of the $\text{Mn}-\text{O}-\text{F}$ anode is comparable or even superior to that of many state-of-the-art anode materials for Li-ion batteries (LIBs) (Table S5, ESI[†]) (the more detailed performance from the CV curves, GCD curves, rate capability and coulombic efficiency, cycling stability and coulombic efficiency of the MnF_2 (1#–9#) and MnF_2 8#-E electrodes can be seen in Fig. S7–S14[†]). The enhanced performance of the $\text{Mn}-\text{O}-\text{F}$ ultrafine nanowires with the hetero oxygen doping and fluorine vacancies provides an important strategy for the promotion of charge storage capability of Li-ion anode materials. Herein, the initial increase in capacity for the MnF_2 8# and MnF_2 8#-E electrodes may be owing to the formation of continuous conductivity networks *via* the conversion, reversible reactions of SEI films and enhanced electroactive sites by the activation of the electrode,³² while the subsequent decrease in capacity for the MnF_2 8#-E electrode may be owing to the decrease in the electroactive sites by the agglomeration of the amorphous ultrafine nanoparticles. The XRD patterns of the electrodes after cycling (Fig. S15A and B, ESI[†]) demonstrate the very faint diffraction peaks of MnF_2 phase compared with the pristine states, and the TEM images of the electrodes after cycling (Fig. S15C–F, ESI[†]) show the presence of numerous ultrafine nanoparticles in the pristine nanocrystals or nanowires, suggesting the formation of a largely amorphous nanostructure for Li-ion storage after the long-term conversions. The CV plots at different sweep rates (0.1, 0.2, and 0.3 mV s⁻¹) of the MnF_2 8# and MnF_2 8#-E electrodes are shown in Fig. 4G and H, and the relationship of $\lg i - \lg v$ ($i = av^b$)³⁶ for MnF_2 8# and MnF_2 8#-E electrodes derived from the CV plots are shown in the insets of

Fig. 4G and H. Note that the slopes of ba and bc are 0.88 and -1.32 for MnF_2 8#, in comparison with that of 0.31 and -0.44 for MnF_2 8#-E, respectively. These results indicate that the kinetics of the MnF_2 8# electrode is dominated by surface-controlled kinetics (*i.e.* pseudocapacitive behavior),³⁷ whereas the MnF_2 8#-E electrode has the typical diffusion-controlled property. Herein, the absolute b values (0.31/0.44) of the MnF_2 8#-E electrode are lower than 0.5, indicating the deviation from the typical planar diffusion characteristics, which may be owing to the highly porous and rough structure of the electrode, leading to the partial spherical diffusion. Moreover, the pseudocapacitive contribution ($k_1 v$)^{36,37} at 0.1–0.3 mV s⁻¹ was estimated to be 73–82% and 31–44% for the MnF_2 8# and MnF_2 8#-E electrodes, respectively (Fig. S16 and S17, ESI[†]).

Electrochemical impedance spectroscopy (EIS) was used to investigate the electrochemical kinetics of the MnF_2 and MnF_2 8#-E electrodes for Li-ion storage. As shown in Fig. S19,[†] the Nyquist plots include a Z' -axis intercept at the superhigh frequency, one semicircle in the high frequency region, a short sloping line in the intermediate frequency region and an oversized curved arc in the low frequency region, which are considered as the internal ohmic resistance (R_s) (including resistances of electrode and electrolyte, and contact resistances between particles and/or between electrode and current collector), charge transfer resistance (R_{ct}) related to the electrochemical kinetics process, Warburg impedance (W) related to the ion diffusion process and the electronic resistance related to the bulk materials, respectively.³² The equivalent circuit model is shown in Fig. S18 (ESI[†]), and the fitting results are listed in Table S6 (ESI[†]), in which the $\text{Mn}-\text{O}-\text{F}$ electrode shows a smaller R_{ct} (37.33 Ω) than the MnF_2 electrodes (43.09–68.34 Ω), implying the faster electrochemical kinetics in the $\text{Mn}-\text{O}-\text{F}$ electrode because of its superior morphology and porosity.

The MnF_2/AC and $\text{MnF}_2\text{-E}/\text{AC}$ LICs were further assembled with MnF_2 8# and MnF_2 8#-E as the anodes and commercial AC as the cathode together with the anodes pre-charged (pre-lithiation) mode at 0.1 A g⁻¹ for 3.5 turns (Fig. S20[†]). Different active mass ratios of the cathode/anode were examined firstly. For the MnF_2/AC LICs, the performance comparison for different mass ratios in Fig. S22[†] makes it clear that MnF_2/AC (1 : 1) is the optimal LIC (the performance of the AC cathode is shown in Fig. S21,[†] and the voltage windows, GCD curves and CV plots of the MnF_2/AC LICs with different mass ratios are shown in Fig. S23 and S24[†]). For the $\text{MnF}_2\text{-E}/\text{AC}$ LICs, the best mass ratio was determined to be 1 : 2 based on the results shown in Fig. S25 and S26.[†] After selecting the optimal mass ratios of the LICs, different voltage windows (0–4 V and 0–4.3 V) were further examined. Fig. 5A–H show the CV plots at 40 mV s⁻¹, GCD curves at 1 A g⁻¹, Ragone behavior and cycling performance at 5 A g⁻¹ of the MnF_2/AC (1 : 1) and $\text{MnF}_2\text{-E}/\text{AC}$ (1 : 2) LICs under the voltages of 0–4 V and 0–4.3 V. The $\text{MnF}_2\text{-E}/\text{AC}$ capacitor exhibited a larger CV area and longer charge and discharge times, indicating its better performance (the CV plots at different sweep speeds and GCD curves under different current densities are shown in Fig. S26–S28[†]). Furthermore, the $\text{MnF}_2\text{-E}/\text{AC}$ LICs exhibited remarkable performances under both 0–4 V (106.8–41.4 W h kg⁻¹/0.33–10.7 kW kg⁻¹, 63.4%



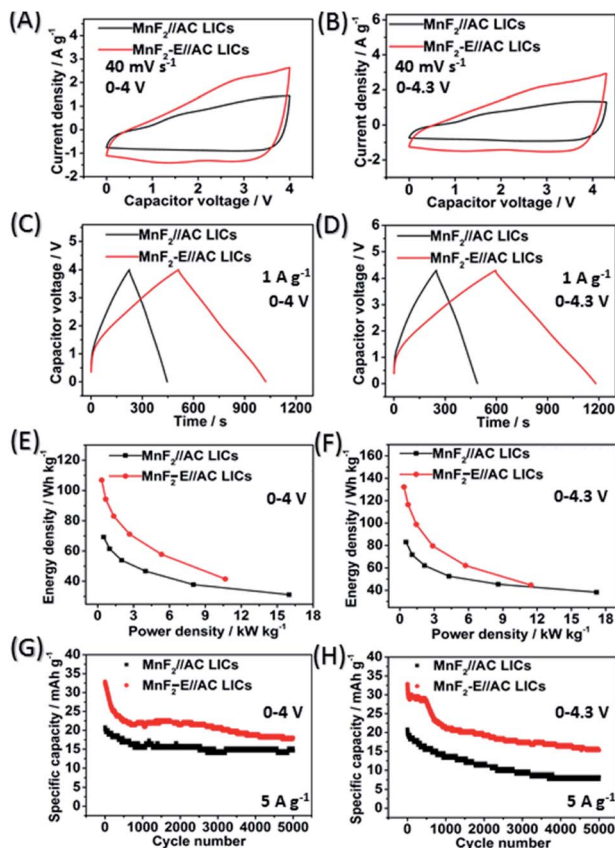


Fig. 5 CV plots at 40 mV s^{-1} (A and B), GCD curves at 1 A g^{-1} (C and D), Ragone plots (E and F) and cycling behavior for 5000 cycles at 5 A g^{-1} (G and H) of the MnF_2 8#/AC and MnF_2 8#-E//AC LICs under the voltages of 0–4 V and 0–4.3 V, respectively.

retention/3000 cycles/ 5 A g^{-1}) and 0–4.3 V ($132.2\text{--}44.6 \text{ Wh kg}^{-1}$ / $0.36\text{--}11.5 \text{ kW kg}^{-1}$, 63.5% retention/1000 cycles/ 5 A g^{-1}) in comparison with the MnF_2 //AC LICs at 0–4 V ($69.2\text{--}31.1 \text{ Wh kg}^{-1}$ / $0.5\text{--}16.0 \text{ kW kg}^{-1}$, 72.5% retention/3000 cycles/ 5 A g^{-1}) and 0–4.3 V ($82.9\text{--}38.2 \text{ Wh kg}^{-1}$ / $0.54\text{--}17.2 \text{ kW kg}^{-1}$, 63.5% retention/1000 cycles/ 5 A g^{-1}) (more details are shown in Table S7†). The remarkable performance of the MnF_2 -E//AC LICs further proves that the double-defective Mn–O–F ultrafine nanowire anode with hetero oxygen doping and intrinsic fluorine vacancy greatly enhances the energy storage performance. Moreover, the MnF_2 -E//AC and MnF_2 //AC LICs exhibited comparable or even better performance than many state-of-the-art LICs systems reported in the literature (Table S8†), demonstrating their promising application in Li-ion storage.

Performance for Na-ion storage

Due to the remarkable Li-ion storage capability of MnF_2 8# and MnF_2 8#-E, they were further investigated for Na-ion storage. In the CV plots of the MnF_2 8# electrode at 0.1 mV s^{-1} (Fig. 6A), there was a big reduction peak at around 1.03 V and a small reduction peak at about 0.16 V , corresponding to the conversion of MnF_2 into Mn/NaF and formation of SEI films in the first discharge, respectively. Moreover, the Na-ion storage capacity in

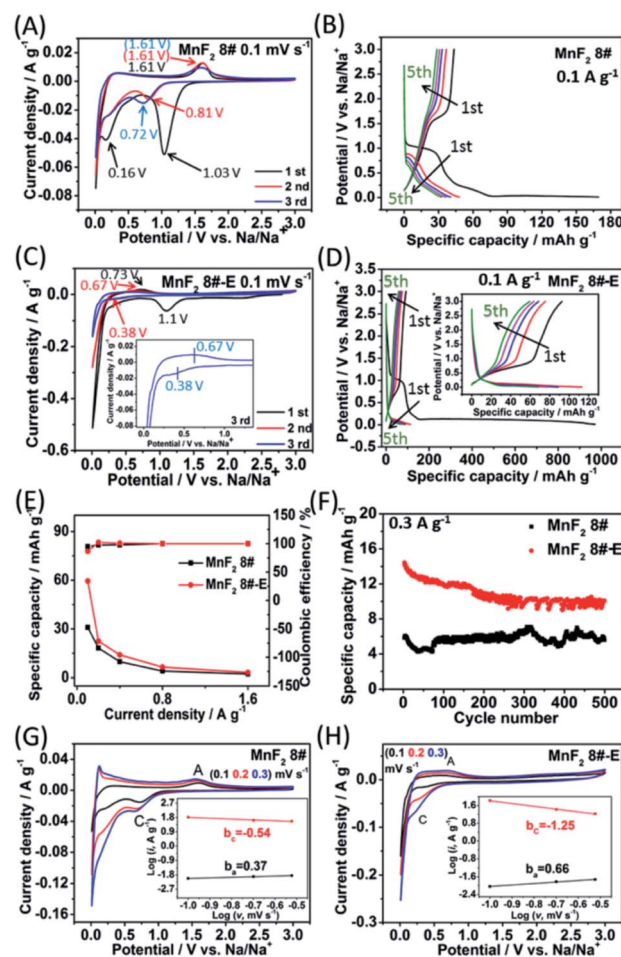


Fig. 6 CV plots for the first three cycles at 0.1 mV s^{-1} (A) and GCD curves for the 5th cycle at 0.1 A g^{-1} (B) of MnF_2 8# electrode, CV plots for the first three cycles at 0.1 mV s^{-1} (C) and GCD curves for the 5th cycle at 0.1 A g^{-1} (D) of the MnF_2 8#-E electrode, specific capacity and coulombic efficiency derived from the respective 5th cycle at $0.1\text{--}1.6 \text{ A g}^{-1}$ of the MnF_2 8# and MnF_2 8#-E electrodes (E), cycling stability for 500 cycles at 0.3 A g^{-1} of the MnF_2 8# and MnF_2 8#-E electrodes (F), and CV plots for the 3rd cycle at $0.1, 0.2, \text{ and } 0.3 \text{ mV s}^{-1}$ of MnF_2 8# and MnF_2 8#-E (insets show the plots of $\lg I$ vs. $\lg v$) (G and H).

the low potential region also originated from a reversible interfacial Na-ion intercalation reaction within the Mn/NaF matrix as Li-ion intercalation.^{31–35} Also, the minor anodic peak at about 1.61 V can be ascribed to the partial conversion of Mn/NaF into MnF_2 . In the second and third laps, the reduction peaks changed lightly from 0.81 V to 0.72 V , and the oxidation peaks were basically maintained at 1.61 V , indicating good reversibility. Fig. 6C shows the CV plots of the MnF_2 8#-E electrode at a sweep speed of 0.1 mV s^{-1} . A large cathode peak can be observed at about 1.1 V and small peak at around 0.4 V in the first round of negative sweep, which can be attributed to the conversion of Mn–O–F into $\text{Mn/NaF/Na}_2\text{O}$ and the formation of SEI films, respectively. Also, the minor anodic peak at 0.73 V during the first positive sweep can be ascribed to the partial conversion of Mn/NaF/ Na_2O into MnF_2/MnO . During the next two turns, the cathode peaks were basically maintained at

0.38 V, while the anode peaks shifted lightly from 0.73 V to 0.67 V, indicating good reversibility. Fig. 6B and D show the GCD curves for the MnF_2 8# and MnF_2 8#-E electrodes in the first five laps of 0.1 A g^{-1} , where the charge/discharge characteristics with the potential platforms coincide with the anodic/cathodic processes with redox peaks in their CV plots. Notably, the potential platforms of the MnF_2 8#-E electrode are obviously lower than that of the MnF_2 8# electrode for Na-ion storage, implying the superiority of the charge storage of the Mn-O-F anode. Fig. 6E and F show the gradient specific capacity and cycle behavior for the MnF_2 8# and MnF_2 8#-E electrodes, where MnF_2 8# provides 31.2 mA h g^{-1} at $0.1\text{--}3.2 \text{ A g}^{-1}$ with 5.4 mA h g^{-1} after 500 cycles at 0.3 A g^{-1} in comparison with the performance of 59.4 mA h g^{-1} at $0.1\text{--}3.2 \text{ A g}^{-1}$ with 11 mA h g^{-1} after 500 cycles at 0.3 A g^{-1} for MnF_2 8#-E (more details are shown in Fig. S29 and S30; Table S4†), which again shows that the double-defective Mn-O-F ultrafine nanowires anodes exhibit a big advantage in Na-ion storage. The XRD patterns of the electrodes after cycling (Fig. S31A and B†) show the slightly smaller diffraction peaks of the MnF_2 phase, and the TEM images of the electrodes after cycling (Fig. S31C-F†) show the presence of a few of ultrafine nanoparticles in the pristine nanocrystals or nanowires, suggesting the formation of partially amorphous nanostructural products for Na-ion storage after the long-term conversions. Fig. 6G and H show the CV plots at different sweep speeds and the relationship between peak current density and sweep rate ($\lg i = b \lg v + a$) for the MnF_2 8# and MnF_2 8#-E electrodes, respectively. The ba and bc values for the MnF_2 8# electrode are 0.37 and -0.54 , respectively, suggesting the typical diffusion-controlled kinetics, and the ba and bc values for the MnF_2 8#-E electrode are 0.66 and -1.25 , respectively, implying the dominant surface-controlled kinetics. Herein, the kinetic behavior of the MnF_2 8# and MnF_2 8#-E electrodes showed obvious differences between Li-ion storage (surface control/diffusion control) and Na-ion storage (diffusion control/surface control), which may be due to the difference in the radius of the Na and Li ions and the surface morphology and structure of the MnF_2 8# and MnF_2 8#-E materials. Moreover, the pseudocapacitive contribution ($k_1 v$)^{36,37} at $0.1\text{--}0.3 \text{ mV s}^{-1}$ was estimated to be 72–82% and 15–24% for the MnF_2 8# and MnF_2 8#-E electrodes (Fig. S32 and S33, ESI†), respectively. The electrochemical kinetics of the MnF_2 8# and Mn-O-F electrodes for Na-ion storage were further analyzed via EIS tests (Fig. S34 and Table S9, ESI†), which show that the Mn-O-F electrode possessed a smaller R_{ct} (106.7Ω) than the MnF_2 electrode (144.8Ω), suggesting the faster electrochemical kinetics of the Mn-O-F electrode owing to its superior morphology and porosity.

The MnF_2 //AC and MnF_2 -E//AC NICs were also assembled with MnF_2 8# and MnF_2 8#-E as anodes and commercial AC as the cathode along with the anodes pre-charged (pre-sodiation) at 0.1 A g^{-1} for 3.5 turns (Fig. S35†). Initially, different active mass ratios of the cathode/anode were examined. The performance comparison for the different mass ratios in Fig. S37† indicate that the MnF_2 //AC (1 : 1) NICs exhibited overall superior behavior (the performance of the AC cathode is shown in Fig. S36,† and the voltage windows, GCD curves and

CV plots of the MnF_2 //AC NICs with different mass ratios are shown in Fig. S38 and S39†). The different potential windows were further examined under the optimal mass ratios of the NICs. Fig. 7A–H show the CV plots at 40 mV s^{-1} , GCD curves at 1 A g^{-1} , Ragone plots, and cycling behavior at 3 A g^{-1} for 800 cycles of the NICs under the voltages of $0\text{--}4 \text{ V}$ and $0\text{--}4.3 \text{ V}$. The MnF_2 8#-E//AC capacitor exhibited a larger CV area and longer charge and discharge times, indicating its better performance (the CV plots at different sweep speeds and GCD curves under different current densities are shown in Fig. S40 and S41,† respectively). The MnF_2 8#-E//AC NICs exhibited a remarkable performance of $42.1\text{--}8.9 \text{ Wh kg}^{-1}$ / $0.5\text{--}8.0 \text{ kW kg}^{-1}$, and 63.1% retention/200 cycles/ 3 A g^{-1} under $0\text{--}4 \text{ V}$ and $55.9\text{--}19.2 \text{ Wh kg}^{-1}$ / $0.48\text{--}7.7 \text{ kW kg}^{-1}$, and 45.3% retention/200 cycles/ 3 A g^{-1} under $0\text{--}4.3 \text{ V}$ in comparison with that of the MnF_2 8#//AC NICs ($16.4\text{--}4.4 \text{ Wh kg}^{-1}$ / $0.5\text{--}8.0 \text{ kW kg}^{-1}$, 50.2% retention/200 cycles/ 3 A g^{-1} / $0\text{--}4 \text{ V}$ and $37\text{--}7.2 \text{ Wh kg}^{-1}$ / $0.54\text{--}8.6 \text{ kW kg}^{-1}$, 66.7% retention/200 cycles/ 3 A g^{-1} / $0\text{--}4.3 \text{ V}$, respectively). Furthermore, the MnF_2 -E//AC NICs and MnF_2 //AC NICs exhibited comparable or even better overall performances than that of many state-of-the-art NICs systems reported in the literature (Table S10†), showing their promising application in Na-ion storage.

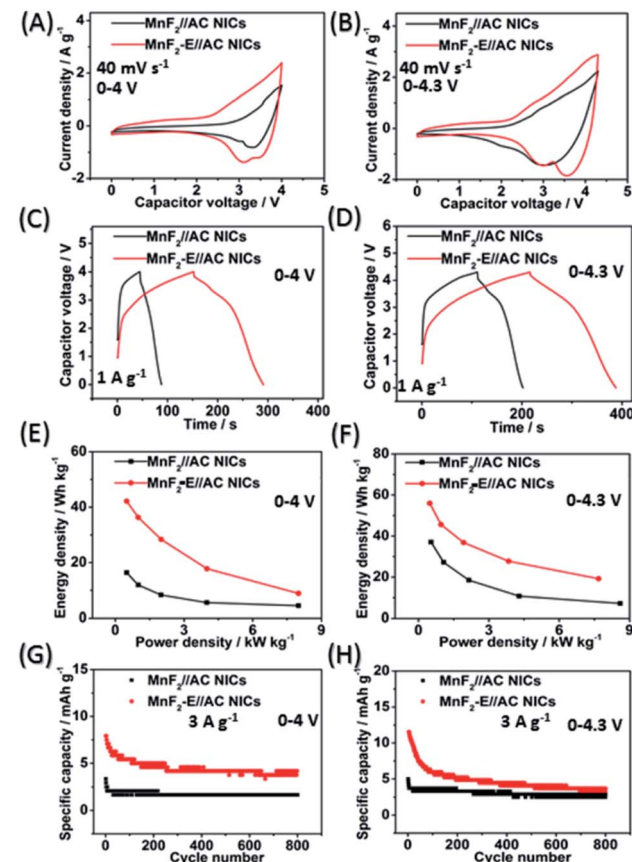


Fig. 7 CV plots at 40 mV s^{-1} (A and B), GCD curves at 1 A g^{-1} (C and D), Ragone plots (E and F), and cycling behavior for 800 cycles at 3 A g^{-1} (G and H) of MnF_2 8#/AC and MnF_2 8#-E//AC NICs under the voltages of $0\text{--}4 \text{ V}$ and $0\text{--}4.3 \text{ V}$, respectively.



Reaction mechanisms for Li/Na-ion storage

Ex situ X-ray photoelectron spectroscopy (XPS) was employed to investigate the possible reaction mechanisms of the MnF_2 and $\text{MnF}_2\text{-E}$ electrodes for both Li-ion and Na-ion storage. Fig. 8 shows the XPS data of the pristine, fully charged (3.0 V) and fully discharged (0.01 V) states of the MnF_2 8# and MnF_2 8#-E anodes

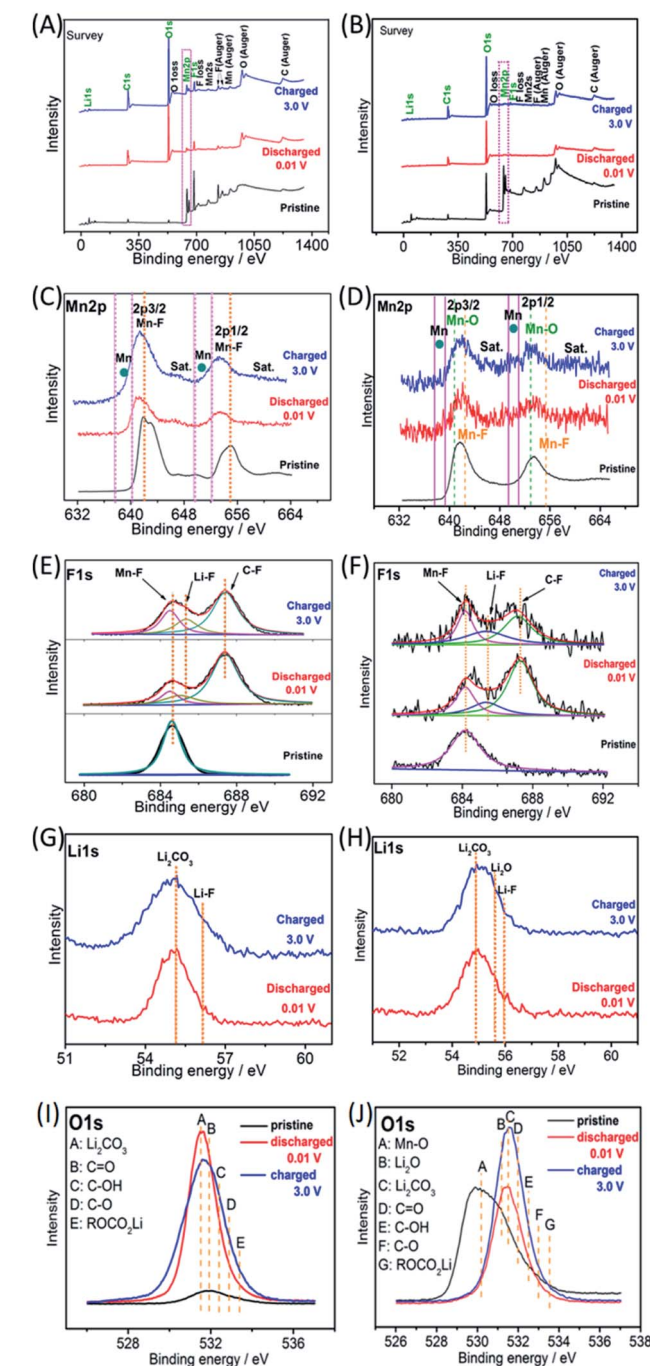


Fig. 8 *Ex situ* XPS spectra of MnF_2 8# and MnF_2 8#-E electrodes in the pristine, charged 3.0 and discharged 0.01 V states during the first discharging/charging cycle at 0.05 A g^{-1} for Li-ion storage: survey (A and B), Mn 2p (C and D), F 1s (E and F), Li 1s (G and H) and O 1s (I and J), respectively.

for the Li-ion storage. The survey scans in Fig. 8A and B, demonstrate the typical F 1s/Mn 2p/O 1s/C 1s in all three states. The much stronger C 1s/O 1s signals of MnF_2 8# in the fully charged/discharged states than the pristine states is largely owing to the formation of SEI films with rich C/O-containing groups.^{32,38,39} For MnF_2 8#-E, the much stronger C 1s signal in the fully charged/discharged states is also owing to the formation of SEI films, while the strong O 1s signals in the fully charged/discharged states originate from both the SEI films and the pristine Mn–O bond. The Mn 2p spectra of MnF_2 8# and MnF_2 8#-E in Fig. 8C and D present the Mn 2p_{3/2} and Mn 2p_{1/2} peaks and corresponding shakeup satellite peaks (Sat.). Notably, the peak positions of Mn 2p showed some negative shifts for the fully discharged and charged states in comparison with that of the pristine states, indicating the partial conversion MnF_2 and Mn–O–F into Mn metal phases during the discharging/charging processes.^{14,31} Moreover, both the Mn–O and Mn–F bonds could be resolved for MnF_2 8#-E in comparison with the sole Mn–F bond for MnF_2 8#. The F 1s spectra in Fig. 8E and F reveal the Mn–F, Li–F and C–F (from polyvinylidene fluoride (PVDF) binder) bonds for the fully discharged/charged states in comparison with the Mn–F bonds in the pristine states. The Li 1s signals in Fig. 8G of the fully discharged and charged states originate from LiF and Li_2CO_3 (from SEI films) during the discharging/charging processes.^{32,38,39} Notably, additional Li_2O signals (except LiF and Li_2CO_3) are present in the Li 1s spectra of MnF_2 8#-E in Fig. 8H, which originated from one of the reaction products of Mn–O–F and Li-ion. Fig. 8I shows the O 1s spectra of MnF_2 8#, which show a weak peak in the original state due to the presence of adsorbed oxygen (mainly H_2O , which can be also detected from the large $-\text{OH}$ absorption peak appearing at 3220 cm^{-1} in the IR spectra of Fig. 1B). The O 1s became sharp under the fully charged/discharged states, which is attributed to the formation of SEI films (mainly including Li_2CO_3 , C=O, C–OH, C–O and ROCO_2Li).^{32,39} Fig. 8J show the O 1s spectra of MnF_2 8#-E, which exhibit a sharp Mn–O bond in its original state, the signals of Li_2CO_3 , C=O, C–OH, C–O and ROCO_2Li in the fully charged/discharged states, suggesting the formation of SEI films, and the additional Li_2O in the fully charged/discharged states, indicating the reaction between Mn–O–F and Li-ion.

The Na-ion storage mechanisms of MnF_2 and $\text{MnF}_2\text{-E}$ were further investigated by *ex situ* XPS measurements of the pristine, fully discharged/charged states in the first cycle. As shown in Fig. 9A and B, the survey scans display the presence of Mn, F, O, Na, C and O species, indicating the reactions between MnF_2 or Mn–O–F electrodes and Na-ion and the formation of SEI films. The Mn 2p spectra of MnF_2 8# in Fig. 9C show the negative shift in the fully charged/discharged states and the presence of Mn, indicating a typical conversion reaction between the MnF_2 and Na-ion. The Mn 2p spectra of MnF_2 8#-E in Fig. 9D demonstrate the presence of Mn, Mn–F and Mn–O bonds, revealing the typical characteristics of the conversion reaction between the Mn–O–F and Na-ion. Fig. 9E and F show the F 1s spectra of MnF_2 8# and MnF_2 8#-E, respectively, where the typical Na–F, Mn–F and C–F (from PVDF) can be resolved,



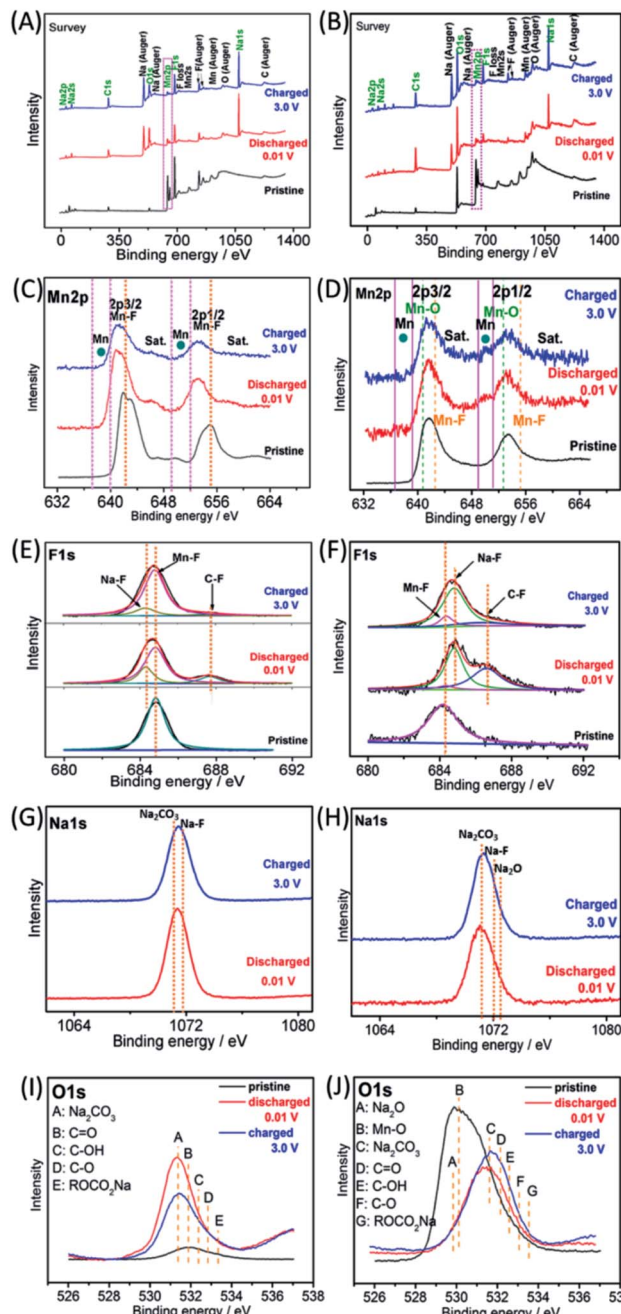
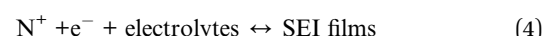
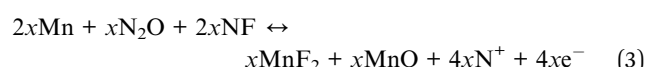
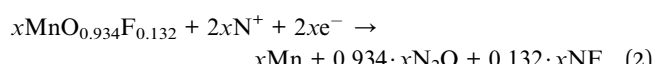
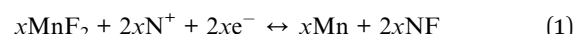


Fig. 9 *Ex situ* XPS spectra of MnF_2 8# and MnF_2 8#-E electrodes in the pristine, charged 3.0 V and discharged 0.01 V states during the first discharging/charging cycle at 0.02 A g^{-1} for Na-ion storage: survey (A and B), Mn 2p (C and D), F 1s (E and F), Na 1s (G and H) and O 1s (I and J), respectively.

which again prove that MnF_2 8# and MnF_2 8#-E exhibit typical conversion reactions. The Na 1s spectra of MnF_2 8# in Fig. 9G indicate the presence of NaF (mainly produced by the conversion reaction) and Na_2CO_3 (from the SEI films). Fig. 9H shows the Na 1s spectra of MnF_2 8#-E, where the Na_2CO_3 bonds indicate the formation of SEI films, and NaF and Na_2O were formed by the feature reactions of Mn-O-F with Na-ion. Fig. 9I shows the O 1s spectra of MnF_2 , where the Na_2CO_3 , C=O, C-OH, C-O and ROCO_2Na bonds are detected. Fig. 9J shows the O 1s spectra of MnF_2 8#-E, where the Na 1s, Mn 2p, F 1s, Na 1s, and O 1s spectra are detected.

and ROCO_2Na bonds in both the fully discharged/charged states are due to the formation of SEI films.^{32,39} Na_2O is also detected in the O 1s spectra of fully discharged/charged MnF_2 8#-E (Fig. 9J), which is derived from the feature reaction of Mn-O-F with Na-ion.

Based on the above considerations, the possible reaction mechanisms of the MnF_2 and Mn-O-F anodes for both Li-ion and Na-ion storage can be expressed as eqn (1)–(6). For MnF_2 , the mechanisms refer to eqn (1), (4) and (5) and for Mn-O-F, the mechanisms refer to eqn (2)–(6). A schematic of the possible processes for the first two discharging/charging cycles is illustrated in Scheme 1.



where $\text{N} = \text{Li, Na}; 0 < x < 1; 0 < y < x$.

Conclusions

In summary, double-defective Mn-O-F ultrafine nanowires with hetero oxygen-doping and intrinsic fluorine-vacancies were introduced as advanced anodes for high-efficiency LICs and NICs. Owing to the unique doping-vacancy double defects, the Mn-O-F anode exhibited a superior performance than the pristine MnF_2 anode, contributing to the superior performance of the Mn-O-F//AC LICs/NICs than the corresponding MnF_2 //AC LICs/NICs. The *ex situ* characterizations and electrochemical methods revealed that the Mn-O-F anode demonstrates typical conversion reaction mechanisms with dominant diffusion-controlled and surface-controlled kinetics for Li/Na-ion storage, respectively. Overall, this work provides new insight into heteroatom-doping and anion vacancy double-defective Mn-O-F ultrafine nanowires anode to improve the Li/Na-ion storage and will have a significant impact on the development of advanced electrochemical energy storage devices.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Thanks for the financial support from National Natural Science Foundation of China (21506182), Natural Science Foundation of Hunan Province (2018JJ2382), Education Department of Hunan

Province (18B077) and Hunan 2011 Collaborative Innovation Center of Chemical Engineering & Technology with Environmental Benignity and Effective Resource Utilization.

Notes and references

- J. M. Tarascon and M. Armand, *Nature*, 2001, **414**, 359–367.
- Z. G. Yang, J. L. Zhang, M. W. KintnerMeyer, X. C. Lu, D. Choi, J. Lemmon and J. Liu, *Chem. Rev.*, 2011, **111**, 3577–3613.
- J. Miller and P. Simon, *Science*, 2008, **321**, 651–652.
- B. Li, J. S. Zheng, H. Y. Zhang, L. M. Jin, D. J. Yang, H. Lv, C. Shen, A. Shellikeri, Y. R. Zheng, R. Q. Gong, J. Zheng and C. M. Zhang, *Adv. Mater.*, 2018, **30**, 1705670.
- H. W. Wang, C. R. Zhu, D. L. Chao, Q. Y. Yan and H. J. Fan, *Adv. Mater.*, 2017, **29**, 1702093.
- J. Ding, W. B. Hu, E. Paek and D. Mitlin, *Chem. Rev.*, 2018, **118**, 6457–6498.
- F. Li and Z. Zhou, *Small*, 2018, **14**, 1702961.
- S. Natarajan, Y.-S. Lee and V. Aravindan, *Chem.-Asian J.*, 2019, **14**, 936–951.
- P. Sennu, N. Arun, S. Madhavi, V. Aravindan and Y.-S. Lee, *J. Power Sources*, 2019, **414**, 96–102.
- P. Pazhamalai, K. Krishnamoorthy, S. Sahoo and S.-J. Kim, *J. Alloys Compd.*, 2018, **765**, 1041–1048.
- G. G. Amatucci, F. Badway, A. D. Pasquier and T. Zheng, *J. Electrochem. Soc.*, 2001, **148**, A930–A939.
- S. Dsoke, B. Fuchs, E. Gucciardi and M. Wohlfahrt-Mehrens, *J. Power Sources*, 2015, **282**, 385–393.
- P. Y. Wang, R. T. Wang, J. W. Lang, X. Zhang, Z. K. Chen and X. B. Yan, *J. Mater. Chem. A*, 2016, **4**, 9760–9766.
- C. F. Liu, C. K. Zhang, H. Q. Song, C. P. Zhang, Y. G. Liu, X. H. Nan and G. Z. Cao, *Nano Energy*, 2016, **22**, 290–300.
- D. P. Dubal, K. Jayaramulu, R. Zboril, R. A. Fischer and P. G. Romero, *J. Mater. Chem. A*, 2018, **6**, 6096–6106.
- R. Ding, Q. Li and H. Y. Wang, *Electrochim. Acta*, 2013, **114**, 726–735.
- N. Yabuuchi, K. Kubota, M. Dahbi and S. Komaba, *Chem. Rev.*, 2014, **114**, 11636–11682.
- E. Lim, C. Jo, M. S. Kim, M. H. Kim, J. Y. Chun, H. Kim, J. Park, K. C. Roh, K. Kang, S. Yoon and J. Lee, *Adv. Funct. Mater.*, 2016, **26**, 3711.
- H. X. Li, J. W. Lang, S. L. Lei, J. T. Chen, K. J. Wang, L. Y. Liu, T. Y. Zhang, W. S. Liu and X. B. Yan, *Adv. Funct. Mater.*, 2018, **28**, 1800757.
- M. L. Kang, Y. Y. Wu, X. Huang, K. Q. Zhou, Z. G. Huang and Z. S. Hong, *J. Mater. Chem. A*, 2018, **6**, 22840–22850.
- B. J. Yang, J. T. Chen, S. L. Lei, R. S. Guo, H. X. Li, S. Q. Shi and X. B. Yan, *Adv. Energy Mater.*, 2018, **8**, 1702409.
- H. L. Wang, D. Mitlin, J. Ding, Z. Li and K. Cui, *J. Mater. Chem. A*, 2016, **4**, 5149–5158.
- Y. L. Huang, Y. X. Zeng, M. H. Yu, P. Liu, Y. X. Tong, F. L. Cheng and X. H. Lu, *Small Methods*, 2018, **2**, 1700230.
- D. F. Yan, Y. X. Li, J. Huo, R. Chen, L. M. Dai and S. Y. Wang, *Adv. Mater.*, 2017, **29**, 1606459.
- S. B. Yang, L. J. Zhi, K. Tang, X. L. Feng, J. Maier and K. Müllen, *Adv. Funct. Mater.*, 2012, **22**, 3634–3640.
- Z. F. Li, C. Bommier, Z. S. Chong, Z. L. Jian, T. W. Surta, X. F. Wang, Z. Y. Xing, J. C. Neuefeind, W. F. Stickle, M. Dolgos, P. A. Greaney and X. L. Ji, *Adv. Energy Mater.*, 2017, **7**, 1602894.
- T. Zhai, X. Lu, Y. Ling, M. Yu, G. Wang, T. Liu, C. Liang, Y. Tong and Y. Li, *Adv. Mater.*, 2014, **26**, 5869.
- Y. C. Wang, T. Zhou, K. Jiang, P. M. Da, Z. Peng, J. Tang, B. Kong, W. B. Cai, Z. Q. Yang and G. F. Zheng, *Adv. Energy Mater.*, 2014, **4**, 1400696.
- Y. W. Liu, H. Cheng, M. J. Lyu, S. J. Fan, Q. H. Liu, W. S. Zhang, Y. D. Zhi, C. M. Wang, C. Xiao, S. Q. Wei, B. J. Ye and Y. Xie, *J. Am. Chem. Soc.*, 2014, **136**, 15670–15675.
- L. Qie, W. M. Chen, Z. H. Wang, Q. G. Shao, X. Li, L. X. Yuan, X. L. Hu, W. X. Zhang and Y. H. Huang, *Adv. Mater.*, 2012, **24**, 2047.
- K. Rui, Z. Y. Wen, Y. Lu, J. Jin and C. Shen, *Adv. Energy Mater.*, 2015, **5**, 1401716.
- D. F. Ying, R. Ding, Y. F. Huang, W. Shi, Q. L. Xu, C. N. Tan, X. J. Sun, P. Gao and E. H. Liu, *J. Mater. Chem. A*, 2019, **7**, 18257–18266.
- P. Balaya, H. Li, L. Kienle and J. Maier, *Adv. Funct. Mater.*, 2003, **13**, 621–625.
- H. Li, G. Richter and J. Maier, *Adv. Mater.*, 2003, **15**, 736–739.
- S. Gruegeon, S. Laruelle, R. Herrera-Urbina, L. Dupont, P. Poizot and J.-M. Tarascon, *J. Electrochem. Soc.*, 2001, **148**, A285–A292.
- J. Wang, J. Polleux, J. Lim and B. Dunn, *J. Phys. Chem. C*, 2007, **111**, 14925–14931.
- V. K. Mariappan, K. Krishnamoorthy, P. Pazhamalai, S. Sahoo, S. S. Nardekar and S.-J. Kim, *Nano Energy*, 2019, **57**, 307–316.
- Q. L. Xu, R. Ding, W. Shi, D. F. Ying, Y. F. Huang, T. Yan, P. Gao, X. J. Sun and E. H. Liu, *J. Mater. Chem. A*, 2019, **7**, 8315–8326.
- W. Shi, R. Ding, Q. L. Xu, T. Yan, Y. X. Huang, C. N. Tan, X. J. Sun, P. Gao and E. H. Liu, *Chem. Commun.*, 2019, **55**, 6739–6742.

